

**COCCULUS CONIUM- cocculus conium pellet**  
**Uriel Pharmacy Inc.**

*Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.*

**Cocculus Conium**

Directions: FOR ORAL USE ONLY.

Dissolve pellets under the tongue 3-4 times daily. Ages 12 and older: 10 pellets. Ages 2-11: 5 pellets. Under age 2: Consult a doctor.

Active Ingredients: Cocculus e fruct. (Cocculus) 4X, Conium mac. (Hemlock) 4X, Ambra grisea (Ambergris) 7X, Petroleum 9X

Inactive Ingredient: Organic sucrose

Use: Temporary relief of dizziness.

**KEEP OUT OF REACH OF CHILDREN.**

Warnings: Claims based on traditional homeopathic practice, not accepted medical evidence. Not FDA evaluated. Contains sugar. Diabetics and persons intolerant of sucrose (sugar): Consult a doctor before use. Do not use if allergic to any ingredient. Consult a doctor before use for serious conditions or if conditions worsen or persist. If pregnant or nursing, consult a doctor before use. Do not use if safety seal is broken or missing.

Questions? Call 866.642.2858 Uriel, East Troy, WI 53120 [www.urielpharmacy.com](http://www.urielpharmacy.com)

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 Lot:

COCCULUS CONIUM			
cocculus conium pellet			
Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:48951-3109
Route of Administration	ORAL		
Active Ingredient/Active Moiety			
Ingredient Name		Basis of Strength	Strength
ANAMIRTA COCCULUS WHOLE (UNII: 8O4P2U3QO2) (ANAMIRTA COCCULUS WHOLE - UNII:8O4P2U3QO2)		ANAMIRTA COCCULUS WHOLE	4 [hp_X]
CONIUM MACULATUM ROOT (UNII: TTE3RU7P5P) (CONIUM MACULATUM ROOT - UNII:TTE3RU7P5P)		CONIUM MACULATUM ROOT	4 [hp_X]
AMBERGRIS (UNII: XTC0D02P6C) (AMBERGRIS - UNII:XTC0D02P6C)		AMBERGRIS	7 [hp_X]

**Inactive Ingredients**

Ingredient Name	Strength
SUCROSE (UNII: C151H8M554)	

**Product Characteristics**

Color	white	Score	no score
Shape	ROUND	Size	3mm
Flavor		Imprint Code	
Contains			

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:48951-3109-2	1350 in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	09/01/2009	

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		09/01/2009	

**Labeler** - Uriel Pharmacy Inc. (043471163)**Establishment**

Name	Address	ID/FEI	Business Operations
Uriel Pharmacy Inc.		043471163	manufacture(48951-3109)